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NEWS 1 Web Page for STN Seminar Schedule - N. America NEWS 2 AUG 10 Time limit for inactive STN sessions doubles to 40

NEWS 2 AUG 10 Time limit for inactive STN sessions doubles to 40 minutes

NEWS 3 AUG 18 COMPENDEX indexing changed for the Corporate Source (CS) field

NEWS 4 AUG 24 ENCOMPLIT/ENCOMPLIT2 reloaded and enhanced

NEWS 5 AUG 24 CA/CAplus enhanced with legal status information for U.S. patents

NEWS 6 SEP 09 50 Millionth Unique Chemical Substance Recorded in CAS REGISTRY

NEWS 7 SEP 11 WPIDS, WPINDEX, and WPIX now include Japanese FTERM thesaurus

NEWS 8 OCT 21 Derwent World Patents Index Coverage of Indian and Taiwanese Content Expanded

NEWS 9 OCT 21 Derwent World Patents Index enhanced with human translated claims for Chinese Applications and Utility Models

NEWS 10 OCT 27 Free display of legal status information in CA/CAplus, USPATFULL, and USPAT2 in the month of November.

NEWS EXPRESS MAY 26 09 CURRENT WINDOWS VERSION IS V8.4, AND CURRENT DISCOVER FILE IS DATED 06 APRIL 2009.

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STRUCTURE FILE UPDATES: 1 NOV 2009 HIGHEST RN 1190833-66-9 DICTIONARY FILE UPDATES: 1 NOV 2009 HIGHEST RN 1190833-66-9

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TSCA INFORMATION NOW CURRENT THROUGH June 26, 2009.

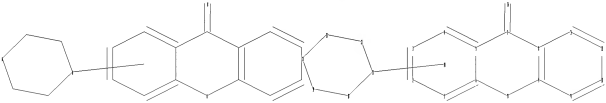
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REGISTRY includes numerically searchable data for experimental and predicted properties as well as tags indicating availability of experimental property data in the original document. For information on property searching in REGISTRY, refer to:

http://www.cas.org/support/stngen/stndoc/properties.html

=>

Uploading C:\Program Files\Stnexp\Queries\QUERIES\10550978.str



Match level :

1:CLASS 2:CLASS 3:CLASS 4:CLASS 5:CLASS 6:CLASS 7:CLASS 8:CLASS 9:CLASS 10:CLASS 11:CLASS 12:CLASS 13:CLASS 14:CLASS 16:CLASS 17:CLASS 18:CLASS 19:CLASS 20:CLASS 21:CLASS 23:CLASS 24:CLASS

=> d L1 HAS NO ANSWERS L1 STR

* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

Structure attributes must be viewed using STN Express query preparation.

=> s 11

SAMPLE SEARCH INITIATED 09:46:36 FILE 'REGISTRY'
SAMPLE SCREEN SEARCH COMPLETED - 62081 TO ITERATE

3.2% PROCESSED 2000 ITERATIONS

3 ANSWERS

INCOMPLETE SEARCH (SYSTEM LIMIT EXCEEDED)

SEARCH TIME: 00.00.01

FULL FILE PROJECTIONS: ONLINE **COMPLETE**

BATCH **COMPLETE**

PROJECTED ITERATIONS: 1226753 TO 1256487 PROJECTED ANSWERS: 1284 TO 2440

L2 3 SEA SSS SAM L1

=> s 11 full

FULL SEARCH INITIATED 09:46:40 FILE 'REGISTRY'
FULL SCREEN SEARCH COMPLETED - 1242745 TO ITERATE

86.9% PROCESSED 1079893 ITERATIONS

2481 ANSWERS

99.0% PROCESSED 1229910 ITERATIONS

2481 ANSWERS

100.0% PROCESSED 1242745 ITERATIONS

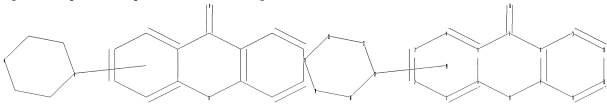
2481 ANSWERS

SEARCH TIME: 00.00.37

L3 2481 SEA SSS FUL L1

=>

Uploading C:\Program Files\Stnexp\Queries\QUERIES\10550978.str



chain nodes :

16

ring nodes :

 $1 \quad 2 \quad 3 \quad 4 \quad 5 \quad 6 \quad 7 \quad 8 \quad 9 \quad 10 \quad 11 \quad 12 \quad 13 \quad 14 \quad 17 \quad 18 \quad 19 \quad 20 \quad 21 \quad 23$

chain bonds :

7-16

ring bonds :

 $1-2 \quad 1-6 \quad 2-3 \quad 3-4 \quad 4-5 \quad 5-6 \quad 5-7 \quad 6-10 \quad 7-8 \quad 8-9 \quad 8-11 \quad 9-10 \quad 9-14 \quad 11-12 \quad 12-13$

13-14 17-23 17-18 18-19 19-20 20-21 21-23

exact/norm bonds :

5-7 6-10 7-8 7-16 9-10 17-23 17-18 18-19 19-20 20-21 21-23normalized bonds :

 $1-2 \quad 1-6 \quad 2-3 \quad 3-4 \quad 4-5 \quad 5-6 \quad 8-9 \quad 8-11 \quad 9-14 \quad 11-12 \quad 12-13 \quad 13-14$

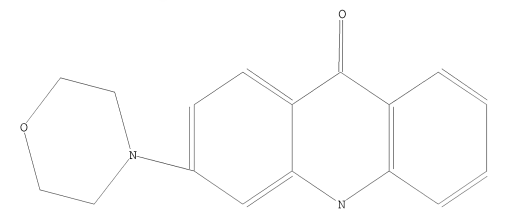
Match level:

1:CLASS 2:CLASS 3:CLASS 4:CLASS 5:CLASS 6:CLASS 7:CLASS 8:CLASS 9:CLASS 10:CLASS 11:CLASS 12:CLASS 13:CLASS 14:CLASS 16:CLASS 17:CLASS 18:CLASS 19:CLASS 20:CLASS 21:CLASS 23:CLASS 24:CLASS

STRUCTURE UPLOADED L4

=> d

L4 HAS NO ANSWERS L4STR



Structure attributes must be viewed using STN Express query preparation.

=> s 14

SAMPLE SEARCH INITIATED 09:51:31 FILE 'REGISTRY' SAMPLE SCREEN SEARCH COMPLETED -146 TO ITERATE

100.0% PROCESSED 146 ITERATIONS 0 ANSWERS

SEARCH TIME: 00.00.01

FULL FILE PROJECTIONS: ONLINE **COMPLETE** BATCH **COMPLETE** PROJECTED ITERATIONS: 2196 TO 3644 PROJECTED ANSWERS: 0 TO

0 SEA SSS SAM L4 L5

=> s 14 full

FULL SEARCH INITIATED 09:51:35 FILE 'REGISTRY' FULL SCREEN SEARCH COMPLETED - 2407 TO ITERATE

100.0% PROCESSED 2407 ITERATIONS 13 ANSWERS SEARCH TIME: 00.00.01

L6 13 SEA SSS FUL L4

=> s 16 and caplus/lc

68915430 CAPLUS/LC L7 12 L6 AND CAPLUS/LC

=> s 16 not 17

L8 1 L6 NOT L7

=> d

```
L8 ANSWER 1 OF 1 REGISTRY COPYRIGHT 2009 ACS on STN
RN 794488-48-5 REGISTRY
ED Entered STN: 08 Dec 2004
9 (1019)-Acridinone, 3-(4-morpholinyl)-1-(phosphonooxy)- (CA INDEX NAME)
MF C17 H17 N2 O6 P
CC CCM
SR CA
```

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

=> fil caplus
COST IN U.S. DOLLARS

FULL ESTIMATED COST

SINCE FILE TOTAL ENTRY SESSION 383.00 383.66

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FILE COVERS 1907 - 3 Nov 2009 VOL 151 ISS 19 FILE LAST UPDATED: 2 Nov 2009 (20091102/ED) REVISED CLASS FIELDS (/NCL) LAST RELOADED: Aug 2009 USPTO MANUAL OF CLASSIFICATIONS THESAURUS ISSUE DATE: Aug 2009

CAplus now includes complete International Patent Classification (IPC) reclassification data for the third quarter of 2009.

CAS Information Use Policies apply and are available at:

http://www.cas.org/legal/infopolicy.html

This file contains CAS Registry Numbers for easy and accurate substance identification.

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=> d his

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FILE 'REGISTRY' ENTERED AT 09:46:08 ON 03 NOV 2009 L1STRUCTURE UPLOADED L2 3 S L1 L3 2481 S L1 FULL STRUCTURE UPLOADED L4L50 S L4 L6 13 S L4 FULL L7 12 S L6 AND CAPLUS/LC 1 S L6 NOT L7 L8

FILE 'CAPLUS' ENTERED AT 09:51:52 ON 03 NOV 2009

=> s 17 L9 2 L7

=> d ibib abs hitstr 1-2

Preparation of xanthenone and acridinone DNA-PK inhibitors as cancer treatment potentiators Halbrook, James W.; Kesicki, Edward A.; Burgess, Laurence Edward; Schlachter, Stephen T.; Eary, TITLE: INVENTOR(S):

Charles

T.; Schiro, Justin G. Icos Corporation, USA PCT Int. Appl., 149 pp. CODEN: PIXXD2 Patent PATENT ASSIGNEE(S): SOURCE:

DOCUMENT TYPE: LANGHAGE English

FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

| | | | | | | | | | | | | LICAT | | | | | | |
|-------|----|-------|------|-----|-----|-----|-----|------|------|-----|----|-------|-------|-----|-----|-----|-------|-----|
| | | 2004 | | | | a 2 | | | 1007 | | | 2004- | | | | | 0040 | |
| | | | | | | | | | | | WO | 2004- | 0504 | 33 | | 2 | 0040. | 313 |
| | WO | 2004 | | | | | | 2005 | | | | | | | | | | |
| | | W: | | | | | | | | | | , BG, | | | | | | |
| | | | CN, | co, | CR, | CU, | CZ, | DE, | DK, | DM, | DZ | , EC, | EE, | EG, | ES, | FI, | GB, | GD, |
| | | | GE, | GH, | GM, | HR, | HU, | ID, | IL, | IN, | IS | , JP, | KE, | KG, | KP, | KR, | KZ, | LC, |
| | | | LK, | LR, | LS, | LT, | LU, | LV, | MA, | MD, | MG | , MK, | MN, | MW, | MX, | MZ, | NA, | NI, |
| | | | NO, | NZ, | OM, | PG, | PH, | PL, | PT, | RO, | RU | , SC, | SD, | SE, | SG, | SK, | SL, | SY, |
| | | | TJ, | TM, | TN, | TR, | TT, | TZ, | UA, | UG, | US | , UZ, | VC, | VN, | YU, | ZA, | ZM, | ZW |
| | | RW: | BW, | GH, | GM, | KE, | LS, | MW, | MZ, | SD, | SL | , SZ, | TZ, | UG, | ZM, | ZW, | AM, | AZ, |
| | | | BY, | KG, | KZ, | MD, | RU, | TJ, | TM, | AT, | BE | , BG, | CH, | CY, | CZ, | DE, | DK, | EE, |
| | | | ES, | FI, | FR, | GB, | GR, | HU, | IE, | IT, | LU | , MC, | NL, | PL, | PT, | RO, | SE, | SI, |
| | | | SK, | TR, | BF, | ВJ, | CF, | CG, | CI, | CM, | GA | , GN, | GΩ, | GW, | ML, | MR, | NE, | SN, |
| | | | TD, | | | | | | | | | | | | | | | |
| | AU | 2004: | 2238 | 66 | | A1 | | 2004 | 1007 | | AU | 2004- | 2238 | 66 | | 2 | 0040 | 319 |
| | CA | 2523 | 178 | | | A1 | | 2004 | 1007 | | CA | 2004- | 2523 | 178 | | 2 | 0040 | 319 |
| | EP | 1660 | 473 | | | A2 | | 2006 | 0531 | | EP | 2004- | 75 78 | 91 | | 2 | 0040 | 319 |
| | | R: | AT, | BE, | CH, | DE, | DK, | ES, | FR, | GB, | GP | , IT, | LI, | LU, | NL, | SE, | MC, | PT, |
| | | | IE, | SI, | FI, | RO, | CY, | TR, | BG, | CZ, | EF | , HU, | PL, | SK | | | | |
| | JP | 2006 | 5236 | 31 | | т | | 2006 | 1019 | | JP | 2006- | 5073 | 73 | | 2 | 0040 | 319 |
| | US | 2007 | 0167 | 441 | | A1 | | 2007 | 0719 | | US | 2006- | 5509 | 78 | | 2 | 0061 | 211 |
| PRIOR | | | | | | | | | | | | 2003- | | | | | | |
| | | | | | | | | | | | WO | 2004- | US84 | 59 | , | w 2 | 0040 | 319 |

MARPAT 141:314155 OTHER SOURCE(S):

ANSWER 1 OF 2 CAPLUS COPYRIGHT 2009 ACS on STN (Continued)

RN 767357-86-8 CAPLUS
CN 3-Acridinecarboxylic acid,
9,10-dihydro-8-hydroxy-6-(4-morpholinyl)-9-oxo, hexyl ester (CA INDEX NAME)

767357-42-6P, 10-Benzyl-1-hydroxy-3-(morpholin-4-yl)-10H-acridin-9-one 767357-70-0P, 10-Benzoyl-1-hydroxy-3-(morpholin-4-yl)-10H-acridin-9-one 767357-71-1P, 1-Hydroxy-10-isobutyryl-3-(morpholin-4-yl)-10H-acridin-9-one 767357-72-2P, 1-Hydroxy-3-(morpholin-4-yl)-10-1(pyridin-4-yl)-10H-acridin-9-one 767357-73-3P, IT

1-Hydroxy-3-(morpholin-4-yl)-10-[(pyridin-3-yl)carbonyl]-10H-acridin-9-one 767357-87-9P, 8-Hydroxy-6-(morpholin-4-yl)-9-oxo-9,10-dihydroacridine-3-carboxylic acid 2-dimethylaminoethyl ester RL: PAC (Pharmacological activity); SPM (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (USES)

(DNA-PK inhibitor; preparation of xanthenone and acridinone DNA-PK inhibitors as cancer treatment potentiators)
767357-42-6 CAPLUS
9(10R)-Acridinone, 1-hydroxy-3-(4-morpholiny1)-10-(phenylmethy1)- (CA

767357-70-0 CAPLUS 9(10H)-Acridinone, 10-benzoyl-1-hydroxy-3-(4-morpholinyl)- (CA INDEX NAME)

ANSWER 1 OF 2 CAPLUS COPYRIGHT 2009 ACS on STN

$$(\mathbb{R}^2)_n - \mathbb{A} = \mathbb{Z}^{\mathbb{Z}^2} \times \mathbb{Z}^{\mathbb{$$

Title compds. I [wherein m = 0-3; n = 0-4; X = 0, S00-2, NRa; Z = independently CRb, N; A = heteroaryl; R1 = independently halo (un) substituted (cyclo)alkyl, heterocyclylalkyl, amino carboxy, phosphoryl, acyl, (hetero)aryl, etc.; R2 = independently halo, CHO, (un) substituted alkyl, (hetero)aryl, carbomyl, carboxy, etc.; R1 = H, (cyclo)alkyl, (hetero)aryl, carboxy, carbamoyl, etc.; Rb = independently H, alkyl, halo, CHO, alkoxy, phosphoryl, amino, carboxy, etc.; and pharmaceutically acceptable salts and prodrugs thereof] were prepared as DNA-dependent protein kinase (DNA-PK) inhibitors. I and their pharmaceutical compns. potentiate cancer treatment by sensitizing cells

(Continued)

an agent that induces DNA lesions. For example, condensation of 1,3-dihydroxy-10H-acridin-9-one with trifluoromethanesulfonic anhydride gave the triflate. Pd-catalyzed substitution of the monoester with morpholine, followed by benzylation provided II. The latter inhibited DNA-PK induced phosphorylation of a p53 peptide substrate with a IC50 of

DNA-PK induced phosphorylation or a processor.

20 nM.

767357-68-97P, 1-Hydroxy-3-(morpholin-4-yl)-10H-acridin-9-one

767357-86-8P, 8-Hydroxy-6-(morpholin-4-yl)-9-oxo-9,10
dihydroacridine-3-carboxylic acid hexyl ester

RL: PRC (Pharmacological activity); RCT (Reactant); SPN (Synthetic

preparation); TRU (Therapeutic use); BIOL (Biological study); PREP

(Preparation); RACT (Reactant or reagent); USES (Uses)

(DNA-PK inhibitors as cancer treatment potentiators)

767357-69-7 CAPLUS

4/1041-Acridinone, 1-hydroxy-3-(4-morpholinyl)- (CA INDEX NAME)

ANSWER 1 OF 2 CAPLUS COPYRIGHT 2009 ACS on STN (Continued)

RN 767357-71-1 CAPLUS CN 9(10H)-Acridinone, 1-hydroxy-10-(2-methyl-1-oxopropyl)-3-(4-morpholinyl)-(CA INDEX NAME)

767357-72-2 CAPLUS 9(10H)-Acridinone, 1-hydroxy-3-(4-morpholiny1)-10-(4-pyridiny1carbony1)-(CA INDEX NAME)

767357-73-3 CAPLUS 9(10H)-Acridinone, 1-hydroxy-3-(4-morpholiny1)-10-(3-pyridinylcarbonyl)-(CA INDEX NAME)

RN 767357-87-9 CAPLUS
CN 3-Acridinecarboxylic acid,
9,10-dihydro-8-hydroxy-6-(4-morpholinyl)-9-oxo, 2-(dimethylamino)ethyl ester (CA INDEX NAME)

767357-89-1, 6-Fluoro-1-hydroxy-3-(morpholin-4-yl)-10H-acridin-9-

one
RL: RCT (Reactant); RACT (Reactant or reagent)
(preparation of xanthenone and acridinone DNA-PK inhibitors as cancer treatment potentiators)
76357-89-1 CAPLOS
9(10H)-Acridinone, 6-fluoro-1-hydroxy-3-(4-morpholinyl)- (CA INDEX NAME)

IT

767357-77-7P, Phosphoric acid dibenzyl ester 3-(morpholin-4-yl)-9-oxo-9,10-dihydroacridin-1-yl ester 767357-81-3P

//o/35/-01-34
//o/36/-01-34
//o/36/-01-34</p

(prodrug, DNA-PK inhibitor; preparation of xanthenone and acridinone

inhibitors as cancer treatment potentiators)

L9 ANSWER 2 OF 2 CAPLUS COPYRIGHT 2009 ACS on STN
ACCESSION NUMBER: 1992:235461 CAPLUS
DOCUMENT NUMBER: 116:235461
CRIGINAL REFERENCE NO: 116:39800h,39801a
Preparation of 9,10-dihydroacridine-9-one derivatives
INVENTOR(S): Butlin, Roger John; Clarvey, Dickson
PATENT ASSIGNEE(S): Imperial Chemical Industries PLC, UK
BUT. Pat. Appl., 33 pp.
DOCUMENT TYPE: Patent
LANGUAGE: Patent
LANGUAGE: English
FAMILY ACC. NUM. COUNT: 1

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

| PATENT NO. | KIND DATE | APPLICATION NO. | DATE |
|------------------------|-----------------|-------------------------|----------|
| | | | |
| EP 471516 | A1 19920219 | EP 1991-307341 | 19910809 |
| R: AT, BE, CH, | DE, DK, ES, FR, | GB, GR, IT, LI, LU, NL, | SE |
| HU 58293 | A2 19920228 | HU 1991-2511 | 19910726 |
| AU 9181422 | A 19920220 | AU 1991-81422 | 19910729 |
| ZA 9105937 | A 19920429 | ZA 1991-5937 | 19910729 |
| CA 2048298 | A1 19920217 | CA 1991-2048298 | 19910801 |
| NO 9103187 | A 19920217 | NO 1991-3187 | 19910815 |
| JP 04257563 | A 19920911 | JP 1991-205027 | 19910815 |
| FI 9103893 | A 19920217 | FI 1991-3893 | 19910816 |
| PRIORITY APPLN. INFO.: | | GB 1990-18044 F | 19900816 |

MARPAT 116:235461 OTHER SOURCE(S):

R SOURCE(S): MARPAT 116:25341
For diagram(s), see printed CA Issue.
Title compds. I (A together with the adjacent vinylene completes a
(substituted) benzene or pyridine ring; R1, R2 = C1-4 alkyl, C1-4 alkoxy;
R3 = H, C1-4 alkyl, C1-4 alkoxy) or a salt, or in vivo hydrolyzable ester
useful as anticancer drugs (no data), are prepared NaH was added to
9-acridinone, followed by 3,5,4-(MeO)2(Me3CSiMe2O)C6H2CH2C1 (preparation
m)

given)

to give the appropriate silylacridinone derivative which was treated with Bu4N+ F- to give after workup I (A completes a benzene ring, R1 = R2 =

Me,

IT 141328-36-1P

IT 141328-36-1P
RL: BAC (Biological activity or effector, except adverse); BSU
(Biological)
study, unclassified); SFN (Synthetic preparation); THU (Therapeutic use);
BIOL (Biological study); PREF (Preparation); USES (Uses)
(preparation of, as anticancer agent)
RN 141328-36-1 CAPLUS
CN 9(10H)-Acridinone, 10-[(4-hydroxy-3,5-dimethoxyphenyl)methyl]-3-(4-morpholinyl)- (CA INDEX NAME)

RN

ANSWER 1 OF 2 CAPLUS COPYRIGHT 2009 ACS on STN (Continued) 767357-77-7 CAPLUS Phosphoric acid, 9,10-dihydro-3-(4-morpholiny1)-9-oxo-1-acridinyl bis(phenylmethyl) ester (CA INDEX NAME)

RN CN (1:2) 767357-81-3 CAPLUS 9(10H)-Acridinone, 3-(4-morpholinyl)-1-(phosphonooxy)-, sodium salt

(CA INDEX NAME)

●2 Na

REFERENCE COUNT:

THERE ARE 10 CITED REFERENCES AVAILABLE FOR

RECORD. ALL CITATIONS AVAILABLE IN THE RE

FORMAT

L9 ANSWER 2 OF 2 CAPLUS COPYRIGHT 2009 ACS on STN (Continued)

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(4 CITINGS)

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DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS)

CA SUBSCRIBER PRICE

SINCE FILE TOTAL
ENTRY SESSION
-1.64
-1.64

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STRUCTURE FILE UPDATES: 1 NOV 2009 HIGHEST RN 1190833-66-9 DICTIONARY FILE UPDATES: 1 NOV 2009 HIGHEST RN 1190833-66-9

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TSCA INFORMATION NOW CURRENT THROUGH June 26, 2009.

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REGISTRY includes numerically searchable data for experimental and predicted properties as well as tags indicating availability of

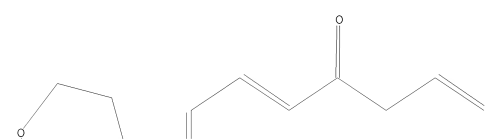
exact/norm bonds : 5-7 6-10 7-8 7-16 9-10 17-23 17-18 18-19 19-20 20-21 21-23 normalized bonds : 1-2 1-6 2-3 3-4 4-5 5-6 8-9 8-11 9-14 11-12 12-13 13-14

Match level :

1:CLASS 2:CLASS 3:CLASS 4:CLASS 5:CLASS 6:CLASS 7:CLASS 8:CLASS 9:CLASS 10:CLASS 11:CLASS 12:CLASS 13:CLASS 14:CLASS 16:CLASS 17:CLASS 18:CLASS 19:CLASS 20:CLASS 21:CLASS 23:CLASS 24:CLASS

L10 STRUCTURE UPLOADED

=> d L10 HAS NO ANSWERS L10 STR



100.0% PROCESSED 40789 ITERATIONS 15 ANSWERS SEARCH TIME: 00.00.01
L12 15 SEA SSS FUL L10

=> d his

L9

(FILE 'HOME' ENTERED AT 09:44:09 ON 03 NOV 2009)

FULL SCREEN SEARCH COMPLETED - 40789 TO ITERATE

FILE 'REGISTRY' ENTERED AT 09:46:08 ON 03 NOV 2009 L1STRUCTURE UPLOADED L23 S L1 L3 2481 S L1 FULL L4STRUCTURE UPLOADED 0 S L4 L5 13 S L4 FULL L6 12 S L6 AND CAPLUS/LC L7 1 S L6 NOT L7 L8 FILE 'CAPLUS' ENTERED AT 09:51:52 ON 03 NOV 2009

2 S L7

FILE 'REGISTRY' ENTERED AT 09:55:16 ON 03 NOV 2009

L10 STRUCTURE UPLOADED L11 0 S L10

L12 15 S L10 FULL

L13 ANSWER 1 OF 2 REGISTRY COPYRIGHT 2009 ACS on STN
RN 383410-09-1 REGISTRY
ED Entered STN: 16 Jan 2002
CN 8H-[1]Benzopyrano[2,3-c]acridine-8,14(13H)-dione,
3-(1,1-dimethylethyl)-6-(4-morpholinyl)- (CA INDEX NAME)
MF C28 H26 N2 O4
SR Chemical Library
Supplier: Ambinter

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

- L13 ANSWER 2 OF 2 REGISTRY COPYRIGHT 2009 ACS on STN NN 382642-26-4 REGISTRY ED Entered STN: 14 Jan 2002 CN 8H-[1]Benzopyrano[2,3-e]acridine-8,14(13H)-dione, 3-methyl-6-(4-morpholinyl)- (CA INDEX NAME) MF C25 H20 N2 O4 SR Chemical Library Supplier: Ambinter

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

(FILE 'HOME' ENTERED AT 09:44:09 ON 03 NOV 2009)

| | FILE | 'REGIS | STRY' | ENTERED | ΑT | 09:46:08 | ON | 03 | NOV | 2009 |
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| L1 | | | STRU | CTURE UP | LOAI | DED | | | | |
| L2 | | 3 | S L1 | | | | | | | |
| L3 | | 2481 | S L1 | FULL | | | | | | |

L4 STRUCTURE UPLOADED

L5 0 S L4 L6 13 S L4 FULL

L7 12 S L6 AND CAPLUS/LC

L9 2 S L7

FILE 'REGISTRY' ENTERED AT 09:55:16 ON 03 NOV 2009 L10 STRUCTURE UPLOADED

L11 0 S L10 L12 15 S L10 FULL

L13 2 S L12 NOT L6

 \Rightarrow s 13 and caplus/lc

68915430 CAPLUS/LC

L14 1391 L3 AND CAPLUS/LC

=> s 13 not 114

L15 1090 L3 NOT L14

 \Rightarrow d 115 1080-1090

L15 ANSWER 1080 OF 1090 REGISTRY COPYRIGHT 2009 ACS on STN
RN 359914-14-0 REGISTRY
ED Entered STN: 03 Oct 2001

4H-Inidazol-4-one, 3,3'-(9-oxo-9H-thioxanthene-3,6-diyl)bis[5-[(4-aminophenyl)methylene]-3,5-dihydro-2-phenyl- (CA INDEX NAME)
F C45 H30 N 60 3 S
SR Chemical Library

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

L15 ANSWER 1082 OF 1090 REGISTRY COPYRIGHT 2009 ACS on STN 87 327080-47-7 REGISTRY Entered STN: 14 Mar 2001
CN Benzimidazo[1',2':1,6]pyrimido[4,5-b]quinolin-14(5H)-one, 5-methyl-7-(2-thienyl)- (CA INDEX NAME)

F C22 H14 N4 o S
Chemical Library Supplier: Ambinter
LC STN Files: CHEMCATS

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

L15 ANSWER 1081 OF 1090 REGISTRY COPYRIGHT 2009 ACS on STN RN 359914-13-9 REGISTRY ED Entered STN: 03 Oct 2001 CN 4H-Imidazol-4-one, 3,3'-(9-oxo-9H-thioxanthene-3,6-diyl)bis[3,5-dihydro-2-phenyl-5-(phenylmethylene)- (CA INDEX NAME) MF C45 H28 N4 03 S SR Chemical Library

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

L15 ANSWER 1083 OF 1090 REGISTRY COPYRIGHT 2009 ACS on STN P. 296246-51-0 REGISTRY COPYRIGHT 2009 ACS on STN P. 2002 Entered STN: 17 Oct 2000 CN 1,3-Dioxolo[4,5-b]acridine-9,10(5H,6H)-dione, 7,8-dihydro-5-hydroxy-7-(4-methoxyphenyl)- (CA INDEX NAME) C21 HI7 N O6 Chemical Library Supplier: Zelinsky Institute of Organic Chemistry LC STN Files: CHEMCATS

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

L15 ANSWER 1084 OF 1090 REGISTRY COPYRIGHT 2009 ACS on STN
RN 112583-94-5 REGISTRY
ED Entered STN: 30 Jan 1988
CN 5H-[1]Benzopyrano[2,3-d]pyrimidin-5-one,
2-[2-(2-methoxyphenyl)-5-oxo-5H-[1]benzopyrano[2,3-b]pyridin-3-yl]- (CA
INDEX NAME)
OTHER CA INDEX NAMES:
CN 5H-[1]Benzopyrano[2,3-b]pyridine, 5H-[1]benzopyrano[2,3-d]pyrimidin-5-one
deriv.
MF C30 H17 N3 O5
CI C0M
SR CA

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

L15 ANSWER 1086 OF 1090 REGISTRY COPYRIGHT 2009 ACS on STN
RN 89491-78-1 REGISTRY
ED Entered STN: 16 Nov 1984
CN 9H-Xanthen-9-one, 2-(ethylthio)-6-(2H-tetrazol-5-yl)-, sodium salt (1:1)
(CA INDEX NAME)
CTHER CA INDEX NAMES:
CN 9H-Xanthen-9-one, 2-(ethylthio)-6-(1H-tetrazol-5-yl)-, sodium salt (9CI)
MF C16 H12 N4 O2 S . Na
CRN (89217-53-8)

L15 ANSWER 1085 OF 1090 REGISTRY COPYRIGHT 2009 ACS on STN
RN 110165-61-2 REGISTRY
ED Entered STN: 05 Sep 1987
CN Hydrazinecarboximidamide, 2-[7-chloro-3,4,9,10-tetrahydro-9-oxo-3-[4-(trifluoromethyl)phenyl]-1(2H)-acridinylidene]-N-hydroxy- (CA INDEX NAME)

) C21 H17 C1 F3 N5 O2 COM CA

NAME) MF (CI (SR (

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

L15 ANSWER 1087 OF 1090 REGISTRY COPYRIGHT 2009 ACS on STN RN 89217-66-3 REGISTRY DE Entered STN: 16 Nov 1984 CN 9H-Xanthene-2-diazonium, 9-oxo-6-(2H-tetrazol-5-yl)- (CA INDEX NAME) CTHER CA ROMEX NAMES: CN 9H-Xanthene-2-diazonium, 9-oxo-6-(1H-tetrazol-5-yl)- (9CI) MF C14 H7 N6 O2 CI CCM

L15 ANSMER 1088 OF 1090 REGISTRY COPYRIGHT 2009 ACS on STN

S9965-52-5 REGISTRY

ED Entered STN: 16 Nov 1984

CN 9H-Thioxanthen-9-one, 6-[5-(1,1,2,2,3,3,3-heptafluoropropyl)-1,3,4-oxadiazol-2-yl]-2-methyl-, 10,10-dioxide (CA INDEX NAME)

OTHER CA INDEX NAMES:

CN 9H-Thioxanthen-9-one, 6-[5-(heptafluoropropyl)-1,3,4-oxadiazol-2-yl]-2-methyl-, 10,10-dioxide (9CI)

OTHER NAMES:

CN HF 374C

MF C19 H9 F7 N2 O4 S

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

- ANSWER 1090 OF 1090 REGISTRY COPYRIGHT 2009 ACS on STN 38187-31-4 REGISTRY
 Entered STN: 16 Nov 1984
 Maphtho[1,2':5,6]pyrano[3,2-e]isoindole-1,3,13(2H)-trione,
 3a,4,13a,13b-tetrahydro-4-(4-methoxyphenyl)-2-phenyl- (CA INDEX NAME)
 C32 H23 N O5
 STN Files: BEILSTEIN*
 (*File contains numerically searchable property data)

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

L15 ANSWER 1089 OF 1090 REGISTRY COPYRIGHT 2009 ACS on STN
RN 59954-02-8 REGISTRY
ED Entered STN: 16 Nov 1984
CN 9H-Thioxanthen-9-one, 3-[5-(1,1,2,2,3,3,3-heptafluoropropyl)-1,3,4oxadlazol-2-yl]-, 10,10-dloxide (CA INDEX NAME)
CTHER CA INDEX NAMES:
CN 9H-Thioxanthen-9-one, 3-[5-(heptafluoropropyl)-1,3,4-oxadiazol-2-yl]-,
10,10-dloxide (GCI)
CTHER NAMES:
CN HF 59C
MF C18 H7 F7 N2 O4 S

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

=> fil caplus COST IN U.S. DOLLARS SINCE FILE TOTAL ENTRY SESSION FULL ESTIMATED COST 219.80 617.74 DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS) SINCE FILE TOTAL ENTRY SESSION CA SUBSCRIBER PRICE -1.640.00

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FILE COVERS 1907 - 3 Nov 2009 VOL 151 ISS 19
FILE LAST UPDATED: 2 Nov 2009 (20091102/ED)
REVISED CLASS FIELDS (/NCL) LAST RELOADED: Aug 2009
USPTO MANUAL OF CLASSIFICATIONS THESAURUS ISSUE DATE: Aug 2009

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L8
             1 S L6 NOT L7
    FILE 'CAPLUS' ENTERED AT 09:51:52 ON 03 NOV 2009
L9
             2 S L7
     FILE 'REGISTRY' ENTERED AT 09:55:16 ON 03 NOV 2009
L10
               STRUCTURE UPLOADED
L11
             0 S L10
            15 S L10 FULL
L12
             2 S L12 NOT L6
L13
          1391 S L3 AND CAPLUS/LC
L14
          1090 S L3 NOT L14
L15
    FILE 'CAPLUS' ENTERED AT 09:58:08 ON 03 NOV 2009
=> s 114
L16
          238 L14
=> s 116 and DNA-PK
       985315 DNA
        21219 DNAS
        988681 DNA
             (DNA OR DNAS)
         27251 PK
         2985 PKS
         29891 PK
                (PK OR PKS)
          1070 DNA-PK
                (DNA(W)PK)
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```
L17 ANSWER 1 OF 1 CAPLUS COPYRIGHT 2009 ACS ON STN
AN 2004:817876 CAPLUS
DN 141:314155
T1 Preparation of xanthenone and acridinone DNA-PK
inhibitors as cancer treatment potentiators
IN Halbrook, James W.; Kesicki, Edward A.; Burgess, Laurence Edward;
Schlachter, Stephen T.; Eary, Charles T.; Schiro, Justin G.
PA Toos Corporation, USA
SO PCT Int. Appl., 149 pp.
CODEN: PIXXD2
DT Patent
LA English
FAN.CNT 1
PATENT NO. KIND DATE APPLICATION NO. DATE

PATENT NO. KIND DATE APPLICATION NO. DATE

PT WO 2004085418 A2 20041007 WO 2004-US8459 20040319
WO 2004085418 A3 20050127
W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BW, BY, BZ, CA, CH,
CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FT, GB, GD,
GE, GH, GH, HR, HU, ID, IL, IN, IS, JF, KE, KG, KF, KR, KZ, LC,
LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, NN, MN, NX, MR, NA, NI,
NO, NZ, CM, FG, PH, PL, FT, RO, RU, SC, SD, SE, SG, SK, SK, LSY,
TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW
RN: BW, GH, GM, KE, LS, MM, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ,
BY, KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK, EE,
ES, FT, FR, GB, RH, LE, TT, LU, NC, NL, PL, PT, RO, FS, SI,
SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN,
TD, TG
AU 200423366 A1 20041007 AU 2004-2523178 20040319
EP 1660473 A2 20060531 EP 2004-757891 20040319
EP 1660473 A2 20060531 EP 2004-5503778 20040319
EP 1660473
```

L16 ANSWER 230 OF 238 CAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 1974:82667 CAPLUS

DOCUMENT NUMBER: 80:13297a,13300a ORIGINAL REFERENCE NO.:

80:1329/a,13300a Heterocyclic-substituted xanthonecarboxylic acid compounds Pfister, Jurg R.; Harrison, Ian T.; Fried, John H. Syntex Corp. Ger. Offen., 45 pp. CODEN: GMXXBX TITLE:

INVENTOR(S): PATENT ASSIGNEE(S):

SOURCE:

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

| | PATENT NO. | KIND | DATE | APPLICATION NO. | DATE |
|-------|-------------------|------|----------|------------------|----------|
| | | | | | |
| | DE 2234258 | A1 | 19731206 | DE 1972-2234258 | 19720712 |
| | US 3835157 | A | 19740910 | US 1972-254232 | 19720517 |
| | NL 7209624 | A | 19731120 | NL 1972-9624 | 19720712 |
| | FR 2184572 | A1 | 19731228 | FR 1972-25346 | 19720712 |
| | JP 49026282 | A | 19740308 | JP 1972-69849 | 19720712 |
| | GB 1394584 | A | 19750521 | GB 1972-32540 | 19720712 |
| | GB 1394585 | A | 19750521 | GB 1974-44679 | 19720712 |
| PRIOR | ITY APPLN. INFO.: | | | US 1972-254232 A | 19720517 |
| | | | | | |

For diagram(s), see printed CA Issue.
Antihistaminic xanthonecarboxylates I (R = 5-methyl-3-furyl,
5-methyl-3-thienyl, 5-methyltetrahydro-3-furyl,
5-methyl-3-thienyl, 5-methyltetrahydro-3-thienyl, 5-methyltetrahydro-3-thienyl, its 1-oxide or 1,1-dioxide,
6-mehyltetrahydropyan-3-yl, 5-methyltetrahydro-thiopyxan-3-yl) were
prepared Thus, 2,4-(Meo2c) 2C6H3Br was treated with p-HOC6H4CH2CH:CH2 and
the 2,4-(Meo2c) 2C6H3CH3CH2-phydrolyzed to the free acid,
cyclized, and esterified to give I (R = CH2CH:CH2). NaIO4 oxidation II

gave

I (R = CH2CHO), which with ClCH2CCMe gave I (R = CH2(CHO)CH2CCMe). Cyclization with acid gave I (R = 5-methyl-3-furyl), and cyclization in the presence of P2S5 yielded I (R = 5-methyl-3-thienyl). S1775-93-0 RL: RCT (Reactant); RACT (Reactant or reagent)

IT

(esterification of)
51775-33-0 CAPUS
9H-Xanthene-2-carboxylic acid, 9-oxo-6-(tetrahydro-3-furanyl)- (CA INDEX

| IT | 51775-80-5P | 51775-81-6P | 51775-82-7P |
|----|------------------|------------------|--------------------|
| | 51775-84-9P | 51775-86-1P | 51775-87-2P |
| | 51775-88-3P | 51775-91-8P | 51775-94-1P |
| | 51775-96-3P | 51775-97-4P | 51775-99-6P |
| | 51776-00-2P | 51823-27-9P | |
| | RL: SPN (Synthet | ic preparation); | PREP (Preparation) |
| | | | |

ANSWER 230 OF 238 CAPLUS COPYRIGHT 2009 ACS on STN (Continued) 51775-86-1 CAPLUS 99H-Xanthene-2-carboxylic acid, 9-oxo-6-(tetrahydro-5-methyl-3-thienyl)-, methyl ester (CA INDEX NAME)

9H-Xanthene-2-carboxylic acid, 9-oxo-6-(tetrahydro-1-oxido-3-thienyl)-, methyl ester (CA INDEX NAME)

RN 51775-88-3 CAPLUS
CN 9H-Xanthene-2-carboxylic acid,
9-oxo-6-(tetrahydro-1,1-dioxido-3-thienyl)-, methyl ester (CA INDEX NAME)

51775-91-8 CAPLUS
9H-Xanthene-2-carboxylic acid,
o-6-(tetrahydro-6-methyl-2H-pyran-3-yl), methyl ester (CA INDEX NAME)

51775-94-1 CAPLUS 9H-Xanthene-2-carboxylic acid, 9-oxo-6-(tetrahydro-3-furanyl)-, methyl

L16 ANSWER 230 OF 238 CAPLUS COPYRIGHT 2009 ACS on STN (Continued)

RN

(prepn. of)
51775-80-5 CAPUS
9H-Kanthene-2-carboxylic acid, 6-(5-methyl-3-furanyl)-9-oxo-, methyl ester

(CA INDEX NAME)

51775-81-6 CAPLUS 9H-Xanthene-2-carboxylic acid, 6-(5-methyl-3-furanyl)-9-oxo- (CA INDEX

 $51775-82-7 \quad {\tt CAPLUS} \\ 9{\tt H-Xanthene-2-carboxylic} \ {\tt acid}, \ 6-(5-{\tt methyl-3-thienyl})-9-{\tt oxo-}, \ {\tt methyl} \\$

(CA INDEX NAME)

51775-84-9 CAPLUS 9H-Xanthene-2-carboxylic acid, 9-oxo-6-(tetrahydro-5-methyl-3-furanyl)-, methyl ester (CA INDEX NAME)

L16 ANSWER 230 OF 238 CAPLUS COPYRIGHT 2009 ACS on STN ester (CA INDEX NAME) (Continued)

51775-96-3 CAPLUS 9H-Xanthene-2-carboxylic acid, 6-(3-furanyl)-9-oxo-, sodium salt (1:1) (CA INDEX NAME)

51775-97-4 CAPLUS RN

9H-Xanthene-2-carboxylic acid, 9-oxo-6-(tetrahydro-3-furanyl)-, ammonium salt (1:1) (CA INDEX NAME)

● NH3

51775-99-6 CAPLUS
9H-Xanthene-2-carboxamide, 9-oxo-6-(3-thienyl)- (CA INDEX NAME)

51776-00-2 CAPLUS

9H-Xanthene-2-carboxylic acid, 6-(3-furanyl)-9-oxo-, compd. with 2-(diethylamino)ethyl 4-aminobenzoate (1:1) (CA INDEX NAME)

CM 1

CRN 51775-95-2 CMF C18 H10 O5

CM

CRN 59-46-1 CMF C13 H20 N2 O2

RN 51823-27-9 CAPLUS CN 9H-Xanthene-2-carboxylic acid, 9-oxo-6-(tetrahydro-5-methyl-2H-thiopyran-3-yl)-, ethyl ester (CA INDEX NAME)

IT

RL: RCT (Reactant); RACT (Reactant or reagent)

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

| PATENT NO. | KIND | DATE | APPLICATION NO. | DATE | |
|--------------------|------|----------|-----------------|-----------|---|
| DE 2234257 | A1 | 19731206 | DE 1972-2234257 | 19720712 | - |
| US 3835158 | A | 19740910 | US 1972=254237 | 1972051 | |
| | | | | 1972031 | |
| NL 7209623 | A | 19731120 | NL 1972-9623 | | _ |
| FR 2184573 | A1 | 19731228 | FR 1972-25348 | 19720712 | |
| AU 7244461 | A | 19740117 | AU 1972-44461 | 19720712 | 2 |
| ZA 7204759 | A | 19740227 | ZA 1972-4759 | 19720712 | 2 |
| JP 49024964 | A | 19740305 | JP 1972-69848 | 19720712 | 2 |
| GB 1393412 | A | 19750507 | GB 1972-32539 | 19720712 | 2 |
| GB 1393413 | A | 19750507 | GB 1974-41751 | 19720712 | 2 |
| GB 1393414 | A | 19750507 | GB 1974-41754 | 19720712 | 2 |
| AT 325039 | В | 19750925 | AT 1972-5990 | 19720712 | 2 |
| AT 325044 | В | 19750925 | AT 1972-325044 | 19720712 | 2 |
| AT 325045 | B | 19750925 | AT 1972-325045 | 19720712 | 2 |
| IL 39888 | A | 19751015 | IL 1972-39888 | 19720712 | 2 |
| SE 387947 | В | 19760920 | SE 1972-9202 | 19720712 | 2 |
| NO 135826 | В | 19770228 | NO 1972-2498 | 19720712 | 2 |
| BE 799580 | A1 | 19731116 | BE 1973-131149 | 19730516 | 6 |
| DK 7406680 | A | 19750602 | DK 1974-6680 | 19741219 | 9 |
| RITY APPLN. INFO.: | | | US 1972-254233 | A 1972051 | 7 |
| | | | | | |

DK 1972-3478 A 19720712

For diagram(s), see printed CA Issue.
Anti-histaminic xanthonecarboxylic acids I (R = tetrahydro-2-furyl, tetrahydro-2-thingly or its 1-oxide or 1,1-dioxide, 2-furyl, 2-thingly, 2,5-dihydro-5-oxo-2-furyl) were prepared Thus, Me xanthene-2-carboxylate was treated with C1(CH2)3COC1, the II (R = C1(CH2)3CO) reduced, and the II (R1 = C1(CH2)3CH2) cyclized with NaH, oxidized with CrO3-pyridine, followed by hydrolysis of the ester group to give I (R = tetrahydro-2-furyl.
51775-59-87
51775-59-89
51775-59-89
51775-70-39
51775-70-39
51775-71-4P
51775-72-5P
51775-73-69
51775-73-69
51775-74-7P
51823-23-7P
51823-34-8P
RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of) (preparation of)
51775-54-3 CAPLUS
9H-Xanthene-2-carboxylic acid, 9-oxo-6-(tetrahydro-2-furanyl)- (CA INDEX NAME)

DK 1972-3478

A 19720712

IT

PR

L16 ANSWER 230 OF 238 CAPLUS COPYRIGHT 2009 ACS on STN (Continued)

(reaction of, with ammonia)

RN 51775-98-5 CAPLUS
CN 9H-Xanthene-2-carboxylic acid, 9-oxo-6-(3-thienyl)- (CA INDEX NAME)

IT

51775-95-2

RL: RCT (Reactant); RACT (Reactant or reagent)
(reaction of, with sodium hydroxide)

51775-95-2

CAPLUS
9H-Xanthene-2-carboxylic acid, 6-(3-furanyl)-9-oxo- (CA INDEX NAME)

L16 ANSWER 231 OF 238 CAPLUS COPYRIGHT 2009 ACS on STN (Continued)

51775-57-6 CAPLUS 9H-Xanthene-2-carboxylic acid, 9-oxo-6-(tetrahydro-2-thienyl)-, methyl ester (CA INDEX NAME)

51775-58-7 CAPLUS 9H-Xanthene-2-carboxylic acid, 9-oxo-6-(tetrahydro-2-thienyl)- (CA INDEX NAME)

51775-59-8 CAPLUS
9H-Xanthene-2-carboxylic acid, 9-oxo-6-(tetrahydro-1-oxido-2-thienyl)-,
methyl ester (CA INDEX NAME)

51775-60-1 CAPLUS
9H-Xanthene-2-carboxylic acid, 9-oxo-6-(tetrahydro-1-oxido-2-thienyl)-(CA INDEX NAME)

51775-66-7 CAPLUS

9H-Xanthene-2-carboxylic acid, 6-(2-furanyl)-9-oxo- (CA INDEX NAME)

(Continued)

51775-70-3 CAPLUS 9H-Xanthene-2-carboxylic acid, 6-(2,5-dihydro-5-oxo-2-furanyl)-9-oxo-

INDEX NAME)

51775-71-4 CAPLUS

2-carboxylic acid, 6-(2-furanyl)-9-oxo-, sodium salt (1:1) 9H-Xanthene-2-ca (CA INDEX NAME)

51775-72-5 CAPLUS

9H-Xanthene-2-carboxylic acid, 9-oxo-6-(tetrahydro-5-oxo-2-furanyl)-, ammonium salt (1:1) (CA INDEX NAME)

L16 ANSWER 231 OF 238 CAPLUS COPYRIGHT 2009 ACS on STN (Continued)

51823-25-7 CAPLUS
9H-Xanthene-2-carboxylic acid,
o-6-(tetrahydro-1,1-dioxido-2-thienyl), methyl ester (CA INDEX NAME)

9H-Xanthene-2-carboxylic acid, 9-oxo-6-(tetrahydro-2-furanyl)-, methyl ester (CA INDEX NAME)

● NH3

51775-73-6 CAPLUS

-2-carboxamide, 9-oxo-6-(2-thienyl)- (CA INDEX NAME)

51775-74-7 CAPLUS
9H-Xanthene-2-carboxylic acid, 6-(2-furanyl)-9-oxo-, compd. with
2-(diethylamino)ethyl 4-aminobenzoate (1:1) (CA INDEX NAME)

CRN 51775-66-7 CMF C18 H10 O5

CM 2

CRN 59-46-1 CMF C13 H20 N2 O2

ACCESSION NUMBER:

DOCUMENT NUMBER:

ACTORPORATE SOURCE:

DOUGENT TYPE:

LANGUAGE:

DOUGENT SOURCE:

AB Reaction of the title compound (I) with RCHO in EtOH containing EtONa gave the stylenedoxy Debay, J. 4 (metaylenedoxy) phenyl, sources and so

the styrene derivs. II [R = Ph, p-Me-OC6H4, 3,4-(methylenedioxy)phenyl, p-Me2NC6H4, p-O2NC6H4]. Reaction of I with P2S5 and HONH2.HCl gave the thione III and 3-methyl(or 2-hydroxy-1-naphthyl)-5-[2-hydroxy-1-naphthyl) (or methyl)]isoxzole, resp. Reaction of I with H2NNH2.H2O-EtOH at 90° gave 3-methyl(or 2-hydroxy-1-naphthyl)-5-[2-hydroxy-1-naphthyl) (or methyl)]pyrazole. Reaction of I with B2NNH2.H2O-EtOH at 90° gave 3-methyl(or 2-hydroxy-1-naphthyl) (or methyl)]pyrazole. Reaction of I with phthalic anhydride, succlinic anhydride, or maleic anhydride in the presence of ZnC12 6 hr at 200-20° gave the diketones IV (Q = 0-C6H4, CH2CH2, CH1CH, resp.).
38187-28-9P 38187-29-0P RL: SPN (Synthetic preparation); FREP (Preparation) (preparation of) 38187-28-9 CAPLUS
9H-Benzo[a]xanthene-10, 11-dicarboxylic acid, 10, 11, 11a, 12-tetrahydro-12-oxo-3-phenyl- (CA INDEX NAME)

IT

RN

38187-29-0 CAPLUS
9H-Benzo[a]xanthene-10,11-dicarboxylic acid,
10,11,11a,12-tetrahydro-9-(4-methoxyphenyl)-12-oxo- (CA INDEX NAME)

OS.CITING REF COUNT: RECORD

THERE ARE 3 CAPLUS RECORDS THAT CITE THIS

(3 CITINGS)

L16 ANSWER 233 OF 238 CAPLUS COPYRIGHT 2009 ACS ON STN ACCESSION NUMBER: 1972:34084 CAPLUS DOCUMENT NUMBER: 76:354084 CAPLUS CRIGINAL REFERENCE NO.: 76:5519a,5522a

TITLE:

AUTHOR(S):

76:15519a,5522a
Acridone studies. VIII. Preparation and properties of monobromo-, nitro-, amino-, and piperidino-10-methylacridones
Hodgeman, D. K. C., Prager, R. H.
Org. Chem. Dep., Univ. Adelaide, Adelaide, Australia
Australian Journal of Chemistry (1972), 25(1), 191-9
CODEN: AJCHAS; ISSN: 0004-9425
Journal CORPORATE SOURCE: SOURCE:

LANGUAGE:

TAGE: English
The preparation and spectral properties of monobromo-, nitro-, amino-,

DOCUMENT TYPE:

piperidino-10-methylacridone are described.
34811-61-5P
RL: SPN (Synthetic preparation); PREP (Preparation)
(preparation of)
34811-61-5 CAPLUS
9(10H)-Acridinone, 10-methyl-3-(1-piperidinyl)- (CA INDEX NAME)

OS.CITING REF COUNT: THERE ARE 6 CAPLUS RECORDS THAT CITE THIS

(6 CITINGS)

L16 ANSWER 234 OF 238 CAPLUS COPYRIGHT 2009 ACS on STN
ACCESSION NUMBER: 1962:462600 CAPLUS
DOCUMENT NUMBER: 57:62600
ORIGINAL REFERENCE NO.: 57:12413h-1,12414a-d
Diels-Alder reaction. Experiments with
2,6-distyryl-y-pyrone and 2-styrylchromones
AUTHOR(S): Aziz, Gamil
CORPORATE SOURCE: Univ. Cairo, Giza, Egypt
SOURCE: CODEN: JOCABH, ISSN: 0022-3263
DOCUMENT TYPE: Journal of Organic Chemistry (1962), 27, 2954-7
CODEN: JOCABH, ISSN: 0022-3263
DOCUMENT TYPE: Journal allale
LANGUAGE: Unavailable
AB Expts. were carried out with 2,6-distyryl-y-pyrone (I) and
2-styrylchromones. 2,6-Dimethylpyrone (I g.) in 15 ml. alc. treated with
1.7 g. BzB, left overnight with 0.43 g. Na in 15 ml. alc., and
crystallized
qave 0.75 g. 1, flakes, m. 168° (alc.). I (I g.) and 1.55 g.
trans-dibenzoylethylene in 50 ml. anisole refluxed 25 hrs. gave 0.25 g.
4a,5,6,7-tetrahydro-5,6-dibenzoyl-7-phenyl-2-styrylchromone (II), m.
242°. Repetition of the reaction using 1 g. I and 3.1 g.
trans-dibenzoyl-ethylene and heating 35 hrs. gave 0.3 g. II. II (0.3 g.)
in 200 ml. alc. refluxed 3 hrs. with 0.78 g. NBZOH.BCl and 1.2 g. NaOAc
gave unchanged II. When the reaction was repeated using PhNHNH2 and
NaOAC
or 2,4 dinitrophenyl-hydrazine, II was still recovered. The following

or 2,4 dinitrophenyl-hydrazine, II was still recovered. The following adducts were formed by the above method (compound formed, g. styryl

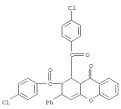
adducts were formed by the above method (compound, addition, g. dienophile used, solvent, ml. solvent, time of heating, solvent of crystallization, m.p. of product, and yield in g. given):

4a, 5, 6, 7-tetrahydro-5, 6-di-p-toluoyl-7-phenyl-2-styrylchromone, 1, 1.76, anisole, 50, 25, alc.-C66h, 266°, 0.25;

4a, 5, 6, 7-tetrahydro-5, 6-dichlorodibenzoyl-7-phenyl-2-styrylchromone, 1, 0.25; 4a,5,6,7-tetrahydro-5,6-dichlorodibenzoyl-7-phenyl-2-styrylchromone, 1, 2.34, anisole, 40, 30, C6H6-ligroine, 248°, 0.24; 7-phenyl-2-styryl-4a,5,6,7-tetrahydrochromone-N-phenyl-5,6-dicarboximide, 0.5, 1.1, xylene, 30, 10, Me2CO, 292°, 0.26. 2-styrylchromone (1 g.) and 1.22 g. trans-p.p'-dichlorodibenzoylethylene, in 40 ml. anisole heated 30 hrs. gave 0.05 g. starting material and 0.15 g. 1,2,3,9a-tetra-hydro-1,2-bis(p-chlorobenzoyl)-9-oxo-3-phenylxanthene, m. 21.5°. The following results were similarly obtained (adduct, styryl compound, g. compound, g. dienophile, time of heating, volume of ole,

styryl compound, g. compound, g. dienophile, time of heating, volume of ole, solvent of crystallization, m.p. of product, and yield in g. given): 5a,6,7,8-tetrahydro-6,7-bis(p-chloro-benzoyl)-4,11-dimethoxy-4-oxo-5H-8-phenylfuro(3,2-b)-xanthene, 2-styrylkhellin, 0.7, 0.6, 30, 2.5, C6H6-ligroine, 230°, 0.21; 5a,6,7,8-tetrahydro-6,7-dibenzoyl-4-methoxy-5-oxo-5H-9-phenylfuro(3,2-b)-xanthene, 2-styrylvisnagin, 1, 0.42, 30, 35, alc.-C6H6, 246°, 0.26; 0.26; 0.26; 0.35,6,7,8-tetrahydro-6,7-bis(p-chlorobenzoyl)-4-methoxy-5-oxo-5H-phenylfuro(3,2-b]-xanthene, 2-styrylvisnagin, 0.58, 0.5, 35, 25, C6H6-ligroine, 258°, 0.25. 96975-47-2P, Xanthen-9-one, 1,2-bis(p-chlorobenzoyl)-1,2,3,9a-tetrahydro-3-phenyl-105003-69-8P, 5H-Furo(3,2-b)xanthen-5-one, 6,7-bis(p-chlorobenzoyl)-5a,6,7,8-tetrahydro-4-methoxy-8-phenyl-1051023-31-7P, 5H-Furo(3,2-b)xanthen-5-one, 6,7-dibenoyl-5a,6,7,8-tetrahydro-4-methoxy-8-phenyl-108040-65-9P, 5H-Furo(3,2-b)xanthen-5-one, 6,7-bis(p-chlorobenzoyl)-5a,6,7,8-tetrahydro-4,11-dimethoxy-8-phenyl-RL: PREP (Preparation) (preparation of)

L16 ANSWER 234 OF 238 CAPLUS COPYRIGHT 2009 ACS ON STN N 96975-47-2 CAPLUS CN 9H-Xanthen-9-one, 1,2-bis(4-chlorobenzoyl)-1,2,3,9a-t-(CA INDEX NAME) 1,2-bis(4-chlorobenzoyl)-1,2,3,9a-tetrahydro-3-phenyl-



RN 105003-69-8 CAPLUS
CN 5H-Furo[3,2-b]xanthen-5-one,
6,7-bis(4-chlorobenzoy1)-5a,6,7,8-tetrahydro4-methoxy-8-pheny1- (CA INDEX NAME)

105123-31-7 CAPLUS 5H-Furo (3,2-b) xanthen-5-one, 7-dibenzoy1-5a,6,7,8-tetrahydro-4-methoxy-8-pheny1- (CA INDEX NAME) L16 ANSWER 234 OF 238 CAPLUS COPYRIGHT 2009 ACS on STN (Continued)

108040-65-9 CAPLUS
5H-Furo[3,2-b]xanthen-5-one,
7-bis(4-chlorobenzoyl)-5a,6,7,8-tetrahydro4,11-dimethoxy-8-phenyl- (CA INDEX NAME)

OS.CITING REF COUNT: 3 THERE ARE 3 CAPLUS RECORDS THAT CITE THIS (3 CITINGS)

116 ANSWER 235 OF 238 CAPLUS COPYRIGHT 2009 ACS on STN (Continued)
6-methyl-3-(o-ethoxyphenyl)-6-methyl-N-(p-ethoxyphenyl), 63,
225-6°, IX, IX, orange; VII, XIV,
3-(3,4-methylenedioxyphenyl)-6-methyl-N-phenyl (XVIII), 54, 215°,
X, XI, brown-red; VII, XV, 3-(3,4-methylenedioxyphenyl)-6-methyl-N-(p-methoxyphenyl), 57, 315, X, brown-red; 2-etryl-7,8-benzochromone
(XIX), XIV, 3-phenyl-N-phenyl-7,8-benzo, 64, 306-8°, IX, XI,
orange; XIX, XV, 3-phenyl-N-(p-methoxyphenyl)-7,8-benzo, 58,
267-9°, X, XII, orange; VIII, XIV,
3-(3,4-methylenedioxyphenyl)-N-[phenyl-7,8-benzo, 72, 288-9°, X, IX,
brown-red; VIII, XV,
3-(3,4-methylenedioxyphenyl)-N-(p-methoxyphenyl)-7,8benzo, 75, 218-9°, X, X, brown-red. XVIII (0.8 g.) in 50 ml. MeOH
with 1 g. NaOH, refluxed for 2 hrs., filtered hot, and acidified with
cold dil. HCl gave 0.35 g. 1,2,3,9a-tetrahydro-9-oxo-3-(3,4-methylenedioxyphenyl)-1,2-xanthenedicarboxylic acid (XX), decomp. 250° (from HOAc). XX is also formed in 75% yield from VII (0.5 g.), 1 g. of maleic acid, and 25 ml. of IX after reflux for 15 hrs. 1082709-14-5p 1082709-32-7p 1087729-31-109 1087729-31-4P RL: SFN (Synthetic preparation); PRP (Properties); PREP (Preparation) (2-Styylchromones in the diene synthesis) 1082709-14-5 CAPLUS

1082709-14-5 CAPLUS
[1]Benzopyrano[3,2-e]isoindole-1,3,11(2H)-trione,
4-(1,3-benzodioxol-5-yl)-3a,4,11a,11b-tetrahydro-2-(4-methoxyphenyl)-8methyl- (CA INDEX NAME)

1082709-32-7 CAPLUS
[1]Benzopyrano[3,2-e]isoindole-1,3,11(2H)-trione,
4-(2-ethoxyphenyl)-3a,4,11a,11b-tetrahydro-8-methyl-2-phenyl- (CA INDEX

L16 ANSWER 235 OF 238 CAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 1957:99120 CAPLUS 51:99120 51:17912f-i,17913a-d DOCUMENT NUMBER: ORIGINAL REFERENCE NO.: 2-Styrylchromones in the diene synthesis Mustafa, Ahmed; Ali, Mohamed Ibrahim Cairo Univ., Egypt Journal of Organic Chemistry (1956), 21, 849-51 TITLE: AUTHOR (S): CORPORATE SOURCE: SOURCE: CODEN: JOCEAH; ISSN: 0022-3263 DOCUMENT TYPE: Journal LANGUAGE: Unavailable
Bo C. C.A. 49, 13234b. Xanthone derivs. are formed by the Diels-Alder reaction from 2-styrylchromones (I) having conjugated double bonds, one which is part of the heterocyclic ring. New I were made by adding alc. NaOEt (from 0.02 g. atom of Na and 20 ml. of absolute EtOH) to 0.02 mole 2-methylchromone and 0.02 mole of an aromatic aldehyde in 40 ml. of EtOH, allowing the mixture to stand at 25° for 24 hrs., filtering off the solid product, washing with cold EtOH (II), and crystallizing from xylene (III). The following chromones were prepared (% yield, m.p., crystallizing solvent, and color in H2SO4 given): 2-(3,4-methylenedioxystyryl) (IV), 209-10°, II, deep red; 2-(p-methoxystyryl)-6-methyl (V), 84, 162-3°, II, red; 2-(o-ethoxystyryl)-6-methyl (VI), 76, 125-6°, III, orange; 2-(3,4-methylenedioxystyryl)-6-methyl (VII), 85, 194-5°, III, deep red; 2-(3,4-methylenedioxystyryl)-7,8-benzo (VIII), 72, 230-1°, III, deep red; I (0,7 g.) and an N-arylmaleimide (1 g.) in 25 ml. of freshly distilled PhOEt (IX), PhNO2 or III were refluxed 10-15 hrs., the resulting solid 1,2,3,9a-tetrahydro-9-oxo-3-aryl-N-aryl'-1,2-xanthenedicarboximides with C6H6, and crystallized from IX, X, Ac2O (XI) or PhOMe (XII). The starting chromone, N-arylmaleimide, 1,2,3,9a-tetrahydro-9-oxo-1,2-xanthenedicarboximide formed, % yield, m.p., reaction solvent, xanthenedicarboximide formed, % yield, m.p., reaction solvent,
tallization
solvent, and color with H2SO4 are listed: 2-styrylchromone (XIII),
N-phenyl (XIV), 3-phenyl-N-phenyl, 75, 252-4°, III, III, crange;
XIII, N-(p-methoxyphenyl) (XV), 3-phenyl-N-(p-methoxyphenyl), 68,
264-6°, III, III, orange; XIII, N-(p-ethoxyphenyl), (XVI),
3-phenyl-N-(p-ethoxyphenyl), 72, 271-3°, III, III, orange; XIII,
N-(2,4-dimethylphenyl), 3-phenyl-N-(2,4-dimethylphenyl), 60,
241-2°, yellow; 2-(p-methoxystyryl)chromone (XVII), XIV,
3-(p-methoxyphenyl)-N-phenyl, 74, 240-1°, III, dioxane-petr. ether
(60-80°), orange-yellow; XVII, XVI,
3-(p-methoxyphenyl)-N-(p-ethoxyphenyl), 65, 214-15°, III, III,
orange; IV, XIV, 3-(3,4-methylenedioxyphenyl)-N-phenyl, 68, above
300°, X, X, red; IV, XV, 3-(3,4-methylenedioxyphenyl)-N-(pmethoxyphenyl), 65, 296-8°, X, X, brown-red; V, XIV,
3-(p-methoxyphenyl)-6-methyl-N-phenyl, 78, 242-3°, IX, IX, yellow;
V, XV, 3-(p-methoxyphenyl)-6-methyl-N-phenyl, 78, 242-3°, IX, IX,
3-(p-methoxyphenyl)-6-methyl-N-(p-ethoxyphenyl), 73,
1(X, orange; VI, XIV, 3-(o-ethoxyphenyl)-6-methyl-N-phenyl, 74,
IX, orange; VI, XIV, yellow; VI, XVI,
3-(3-5°, IX, IX, yellow; VI, XVI, crystallization

L16 ANSWER 235 OF 238 CAPLUS COPYRIGHT 2009 ACS on STN (Continued)

1087729-11-0 CAPLUS [1]Benzopyrano[3,2-e]isoindole-1,3,11(2H)-trione, 3a,4,11a,11b-tetrahydro-4-(4-methoxyphenyl)-8-methyl-2-phenyl- (CA INDEX

1087729-13-2 CAPLUS

1001/23-13-2 CAFEDS [1]Benzopyrano[3, 2-e]isoindole-1,3,11(2H)-trione, 2-(4-ethoxyphenyl)-3a,4,11a,11b-tetrahydro-4-(4-methoxyphenyl)-8-methyl-(CA INDEX NAME)

1087729-23-4 CAPLUS
Naphtho[1',2':5,6]pyrano[3,2-e]isoindole-1,3,13(2H)-trione,
2-(4-ethoxypheny1)-3a,4,13a,13b-tetrahydro-4-(4-methoxypheny1)- (CA) NAME)

1087729-31-4 CAPLUS [1] Benzopyrano[3,2-e]isoindole-1,3,11(2H)-trione,
4-(2-ethoxyphenyl)-2-(4-ethoxyphenyl)-3a,4,11a,11b-tetrahydro-8-methyl-(CA INDEX NAME)

103166-40-1F, Xanthene-1,2-dicarboximide, N-(p-ethoxyphenyl)-1,2,3,9a-tetrahydro-9-oxo-3-phenyl-103271-61-0P, Xanthene-1,2-dicarboximide, IT

L16 ANSWER 235 OF 238 CAPLUS COPYRIGHT 2009 ACS on STN (Continued)

103329-11-9 CAPLUS [1]Benzopyrano[3,2-e]isoindole-1,3,11(2H)-trione,
2-(4-ethoxyphenyl)-3a,4,11a,11b-tetrahydro-4-(4-methoxyphenyl)- (CA CN NAME)

103329-12-0 CAPLUS
[1]Benzopyrano[3,2-e]isoindole-1,3,11(2H)-trione,
3a,4,11a,11b-tetrahydro-2,4-bis(4-methoxyphenyl)-9-methyl- (CA INDEX NAME)

(prepn. of)
103166-40-1 CAPLUS
[1]Benzopyano[3,2-e]isoindole-1,3,11(2H)-trione,
2-(4-ethoxyphenyl)-3a,4,11a,11b-tetrahydro-4-phenyl- (CA INDEX NAME)

103271-61-0 CAPLUS [1]Benzopyrano[3,2-e]isoindole-1,3,11(2H)-trione, 2-(4-ethoxyphenyl)-3a,4,11a,11b-tetrahydro-4-(4-methoxyphenyl)-9-methyl-(CA INDEX NAME)

L16 ANSWER 235 OF 238 CAPLUS COPYRIGHT 2009 ACS on STN (Continued)

114585-07-8 CAPLUS
1H-Vanthene-1,2-dicarboxylic acid,
3-(1,3-benzodioxol-5-yl)-2,3,9,9a-tetrahydro-7-methyl-9-oxo- (CA INDEX

115099-41-7 CAPLUS [1]Benzopyrano[3,2-e]isoindole-1,3,11(2H)-trione,
2-(2,4-dimethylphenyl)-3a,4,11a,11b-tetrahydro-4-phenyl- (CA INDEX NAME)

115099-43-9 CAPLUS
[1]Benzopyrano[3,2-e]isoindole-1,3,11(2H)-trione,
3a,4,11a,1lb-tetrahydro-4-(4-methoxyphenyl)-9-methyl-2-phenyl- (CA INDEX NAME)

115829-14-6 CAPLUS
[1]Benzopyrano[3,2-e]isoindole-1,3,11(2H)-trione,
3a,4,11a,11b-tetrahydro-4-(4-methoxypheny1)-2-pheny1- (CA INDEX NAME)

115829-22-6 CAPLUS
[1]Benzopyrano[3,2-e]isoindole-1,3,11(2H)-trione,
3a,4,1la,1lb-tetrahydro-2-(4-methoxyphenyl)-4-phenyl- (CA INDEX NAME)

115918-34-8 CAPLUS

L16 ANSWER 235 OF 238 CAPLUS COPYRIGHT 2009 ACS on STN (Continued)

121656-69-7 CAPLUS
[1]Benzopyrano[3,2-e]isoindole-1,3,11(2H)-trione,
4-(1,3-benzodioxol-5-yl)-3a,4,11a,11b-tetrahydro-2-(4-methoxyphenyl)-9-methyl- (CA INDEX NAME)

121677-99-4 CAPLUS
[1]Benzopyrano[3,2-e]isoindole-1,3,11(2H)-trione,
4-(1,3-benzodioxol-5-yl)-3a,4,1la,1lb-tetrahydro-9-methyl-2-phenylINDEX NAME) (CA

121678-00-0 CAPLUS
[1]Benzopyrano[3,2-e]isoindole-1,3,11(2H)-trione,
4-(1,3-benzodioxol-5-yl)-3a,4,1la,1lb-tetrahydro-2-(4-methoxyphenyl)-(CA

NAME)

116031-51-7 CAPLUS
[1]Benzopyrano[3,2-e]isoindole-1,3,11(2H)-trione,
4-(2-ethoxyphenyl)-2-(4-ethoxyphenyl)-3a,4,1la,1lb-tetrahydro-9-methyl-(CA INDEX NAME)

117900-61-5 CAPLUS
Naphtho[2',1':5,6]pyrano[3,2-e]isoindole-1,3,13(2H)-trione,
4-(1,3-benzodioxol-5-yl)-3a,4,13a,13b-tetrahydro-2-(4-methoxyphenyl)-

(CA INDEX NAME)

ANSWER 235 OF 238 CAPLUS COPYRIGHT 2009 ACS on STN INDEX NAME)

122117-97-9 CAPLUS
Naphtho[2',1':5,6]pyrano[3,2-e]isoindole-1,3,13(2H)-trione,
3a,4,13a,13b-tetrahydro-2-(4-methoxyphenyl)-4-phenyl- (CA INDEX NAME)

860178-85-4 CAPLUS
[1]Benzopyrano[3,2-e]isoindole-1,3,11(2H)-trione,
3a,4,11a,11b-tetrahydro-4-(4-methoxyphenyl)-4-phenyl- (CA INDEX NAME)

OS.CITING REF COUNT: RECORD

THERE ARE 6 CAPLUS RECORDS THAT CITE THIS

L16 ANSWER 236 OF 238 CAPLUS COPYRIGHT 2009 ACS on STN (Continued) unchanged succinic anhydride (3:5 g.) filtered of, and the filtrate

unchanged succinic anhydride (3:5 g.) filtered of, and the filtrate d.
gave 0.8 g. unchanged II. 2-Methyl-1,4-\(\alpha\)-naphthopyrone (XI) (1 g.)
and 1 g. P2S5 refluxed 2 hrs. in 30 cc. dry C6H6 on a steam bath and
filtered, the residue extd. with boiling C6H6, and the combined filtrates
evapd. to dryness gave 80% 2-methyl-4-thio-\(\alpha\)-naphthopyrone (XII),
violet-red needles, m. 162° (from EtOH), yellow with green
fluorescence in concd. H2S04. XII (1 g.) and 0.5 g. B2H in 20 cc. abs.
EtOH contg. 6 drops piperidine refluxed 6 hrs. and filtered gave 0.8 g.
2-styryl-4-thio-\(\alpha\)-naphthopyrone (XIII), violet crystals, m.
197° (from C6H6), orange in concd. H2S04. XII (1 g.) and 0.6 g.
p-MeOC6H4CHO refluxed gave 0.9 g. p-MeOC6H4CH:CH analog of XIII,
violet
crystals, m. 208° (from C6H6), orange in concd. H2S04. XI (0.1 g.)
in 1 cc. pyridine refluxed 4 hrs. with 0.12 g. NH2OH.HCl in 0.5 cc. H2O,
cooled, acidified with dil. AcOH, and filtered gave 85% 2-[5(or
3)-methyl-3(or 5)-isoxazolyl]-1-naphthol (XIV), yellowish crystals, m.
181° (from C6H6), gave a violet color with alc. FeC13. XIV in 10%
aq. NaOH refluxed 1 hr., cooled, and acidified with dil. HCl gave
unchanged XIV. XIV (0.5 g.) in 10 cc. 10% aq. NaOH shaken 15 min. with
0.5 g. BzCl yielded 0.6 g. Bz deriv., m. 126° (from aq. EtOH). I
(1 g.) in 10 cc. EtOH warmed 15 min. with 50. 50% N2H4.H2O in 10 cc.

warm EtOH, cooled, dild. with H2O, and filtered gave 2-[5(or 3)-methyl-3(or 5)-pyrazolyl]-l-naphthol (XV), colorless leaflets, m. 171°, it gave a deep green color with alc. FeCl3; di-Bz deriv., colorless crystals, m. 144-5° (from aq. EtOH), yellow in concd. ESO4. PhNHNH2.HCl (0.7 g.) in 3 cc. H2O and 1 g. I in 10 cc. pyridine refluxed 4 hrs., cooled, and acidified with dil. AcOH gave 85% l-Ph deriv. (XVI) of XV, almost colorless crystals, m. 143°; it gave a violet color with alc. FeCl3. Il gave similarly the 5(or 3)-styryl analog of XVI, almost colorless needles, m. 223° (decompn.); it gave a violet color with alc. FeCl3.

colnless needles, m. 223° (decompn.); it gave a violet collable. FeCl3.

alc. FeCl3.

122447-99-8P, 7H-Benzo[c]xanthene-8,9-dicarboxylic anhydride,
7a,8,9,10-tetrahydro-10-(p-methoxyphenyl)-7-oxo122324-91-9P,
7H-Benzo[c]xanthene-8,9-dicarboxylic acid,
7a,8,9,10-tetrahydro-10-(p-methoxyphenyl)-7-oxo124483-76-7P,
7H-Benzo[c]xanthene-8,9-dicarboxylic acid,
7a,8,9,10-tetrahydro-10-(p-methoxyphenyl)-7-oxo-, dimethyl ester
RL: FREP (Freparation)
(preparation of)
122447-99-8 CAPUS
3H-Benzo[h]furo[3,4-a]xanthene-1,3,13-trione,
3a,4,13a,13b-tetrahydro-4-(4-methoxyphenyl)- (CA INDEX NAME)

122924-91-8 CAPLUS
7H-Benzo[c]xanthene-8,9-dicarboxylic acid,
7a,8,9,10-tetrahydro-10-(4-methoxyphenyl)-7-oxo- (CA INDEX NAME)

L16 ANSWER 236 OF 238 CAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 1957:9351 CAPLUS

DOCUMENT NUMBER: ORIGINAL REFERENCE NO.: 51:9351 51:1953b-i,1954a-b TITLE:

2-Methyl-1,4- α -naphthopyrone and related substances

AUTHOR(S): Schonberg, Alexander; Fateen, Abd El Kader; Sammour, Abd El Maged Amine

CORPORATE SOURCE:

And El Magdet Amilie Cairo Univ., Egypt Journal of the American Chemical Society (1956), 78, 4689-92

CODEN: JACSAT; ISSN: 0002-7863

DOCUMENT TYPE: Journal Unavailable OTHER SOURCE(S):

ASSEL STATE EtOH

EtOH

treated at room temperature with 1 mole equivalent NaOEt in EtOH, the
mixture treated

with 1 mole equivalent of the appropriate aldehyde, and the yellow
condensation product filtered off and recrystd. gave the corresponding
2-(2-arylstyryl)-1,4-a-naphthopyrone (II); addnl. II was obtained,
in general, by dilution of the mother liquors with H2O; the II was
precipitated in
some cases only upon dilution with H2O. In this manner were prepared the
following II (aryl group, g. weight of I, g. weight of aldehyde used,
m.D.,

color of II, g. yield, and color with H2SO4 given): o-ClC6H4, 1, 0.7,228° (from EtOH or xylene), light yellow, 0.7, yellow (difficultly soluble in C6H6 or ligroine, b. 100-20°); p-O2NC6H4, 1, 0.6, 274-5° (from dioxame or xylene), deep yellow, 0.9, orange (difficultly soluble in C6H6, ligroine, or EtOH); 3,4-(EtO)2C6H3, 1, 0.9,168° (from ligroine), yellow, 0.5, orange (easily soluble in EtOH or C6H6); PhCH-CH3, 1, 0.6, 169° (from aqueous EtOH), yellow-orange, 0.4, deep orange (easily soluble in C6H6 or EtOH, difficultly soluble in ligroine); 3,4-(EEO2)C6H3, 1, 0.7, 232° (from ligroine or EtOH), yellow, 0.6, orange (soluble in C6H6). 2-Styryl-1,4-4-naphthopyrone (III) (2 g.) in 40 cc. 20% aqueous NaOH refluxed 20 hrs. and cooled, the filtrate acidified with H2SO4, washed with H2O, shaken with aqueous 33, Na2CO3,

03, and filtered, and the residue recrystd. from EtOH gave 0.6 g. 2-AcC10H6OH (IV); the filtrate acidified with H2SO4 deposited 0.4 g. FhCH:CBC02H (V), m. 133°. II(0.6 g.) and 5 g. Na in 30 cc. absolute EtOH refluxed 30 hrs. gave 0.6 g. V and 0.8 g. IV. III (g.) and 10 equivs. maleic anhydride (VI) in 30 cc. xylene refluxed 15 hrs., concentrated, and

70% adduct (VII) (R = Ph), almost colorless crystals, m. 279° (from xylene), pale yellow in concentrated H2SO4. p-MeOC6H4CH:CH analog) of III cooled (VIII)

(1) of III (1 g.) and 10 equivs. VI refluxed 15 hrs. in xylene gave 70% VII (R = p-MeOC6H4) (IX), m. 286° (from xylene), yellow in concentrated H2SO4. (0.5 g.) refluxed 50 min. with 1.5 g. NaOH in 18 cc. MeOH, the residue decomposed with HCl, the product dissolved in absolute MeOH, treated 2

dry HCl, allowed to stand overnight, and evaporated, and the residue recrystd.

recrysta.

from MeOH gave the di-Me ester (X) of the corresponding diacid, m.
199°, yellow in concentrated H2SO4. II (1 g.) and 4 g. succinic
anhydride in 30 cc. dry xylene refluxed 15 hrs., concentrated, and
cooled, the

L16 ANSWER 236 OF 238 CAPLUS COPYRIGHT 2009 ACS on STN (Continued)

124483-76-7 CAPLUS
7H-Benzo[c] kanthene-8,9-dicarboxylic acid,
7a,8,9,10-tetrahydro-10-(4-methoxyphenyl)-7-oxo-, 8,9-dimethyl ester (CA
INDEX NAME)

OS.CITING REF COUNT: THERE ARE 3 CAPLUS RECORDS THAT CITE THIS (3 CITINGS)

L16 ANSWER 237 OF 238 CAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 1955:69083 CAPLUS

DOCUMENT NUMBER: 49:69083

ORIGINAL REFERENCE NO.: 49:13234a-q

49:13234a-q Diels-Alder reaction. II. Experiments with 2-styrylchromones. On the nature of the dimer of 1,3-diphenylisobenzofuran Schonberg, Alexander; Mustafa, Ahmed; Aziz, Gamil Cairo Univ., Egypt Journal of the American Chemical Society (1954), 76, 4576-7 TITLE:

AUTHOR(S): CORPORATE SOURCE:

SOURCE:

CODEN: JACSAT: ISSN: 0002-7863

DOCUMENT TYPE: Journal Unavailable

DOCUMENT TYPE:

LANGUAGE: Unavailable
GI For diagram(s), see printed CA Issue.
AB Xanthone derivs. may be obtained from 2-styrylchromones by Diels-Alder reactions. It seems possible that the dimer of 1,3-diphenylisobenzofuran
(I) is a Diels-Alder adduct having the formula II. 2-Styrylchromone

(1 g.) and 4 g. maleic anhydride in 30 cc. dry xylene refluxed 12 h., the mixture concentrated and cooled, and the crystalline deposit filtered washed with hot EtOH, and recrystd. from xylene yielded about 0.65 g. 1,2,3,9a-tetrahydro-9-oxo-3-phenyl-1,2-xanthenedicarboxylic anhydride (IV), m. 246°. 4-Meo derivative of III (0.8 g). gave similarly during 0.5 h. heating in 30 cc. xylene 0.35 g. 3-p-MeoC6H4 analog of IV, m. 260°. Kellin (0.5 g). and 0.25 g. BzH in 10 cc. absolute EtOH treated with cooling with a cooled solution of 0.05 g. Na in 5 cc. absolute, the

mixture warmed slightly, the solution let stand overnight at room

temperature, and
the yellow deposit filtered off, washed with a little EtOH, and recrystd.
from EtOH gave about 0.42 g. 2-styrylkhellin (V), deep yellow needles, m.
196°, easily soluble in warm C6H6, dissolved in concentrated H2S04 with a
red-brown color. Visnaqin and BzH gave similarly about 80%
2-styrylvisnaqin (VI), almost colorless crystals, m. 176° (brownish
melt), dissolved in concentrated H2S04 with an orange color. VI (1.0

g. gave with maleic anhydride during 10 min. in 20 cc. xylene 0.4 g. 5a,6,7,8-tetrahydro-4-methoxy-5-oxo-5H-furo(3,2-b)xanthene-6,7-dicarboxylic anhydride (VII), m. 256° (decomposition) (from Me2CO). (1.0 g.) gave similarly in 25 cc. xylene 0.3 g. 11-MeO derivative of

VII, m. 256° (decomposition) (from dioxane). IV (0.35 g.) refluxed with NaOH in MeOH, the solid product filtered off and decomposed with HCl, and the precipitate

upitate
recrystd. from absolute MeOH gave about 0.21 g.
3,9-dihydro-9-oxo-3-phenyl-1,2-xanthenedicarboxylic acid monohydrate
(VII), m. about 258° (decomposition), soluble in aqueous NAHCO3. VI. treate

with absolute MeOH and dry HCl gave the di-Me ester, colorless crystals,

66-8°. VII (0.25 g.) refluxed with Ac2O gave IV. 2-Styryl-3-methylchromone (0.35 g.) and 2 g. maleic anhydride in 20 cc. ECOPh refluxed 24 h. gave only 0.24 g. recovered starting material.

1089699-71-7P RL: SPN (Synthetic preparation); PRP (Properties); PREP (Preparation) (Diels-Alder reaction. II. Experiments with 2-styrylchromones. On the nature of the dimer of 1,3-diphenylisobenzofuran) 1089699-71-7 CAPLUS

RN

IT

L16 ANSWER 237 OF 238 CAPLUS COPYRIGHT 2009 ACS on STN (Continued)

CAPLUS 1,2-dicarboxylic acid, 2,3,9,9a-tetrahydro-9-oxo-3-phenyl-1H-Xanthene-1,2 (CA INDEX NAME)

CO2H

OS.CITING REF COUNT:

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(4 CITINGS)

L16 ANSWER 237 OF 238 CAPLUS COPYRIGHT 2009 ACS on STN (Cont CN 3H-Xanthene-1,2-dicarboxylic acid, 4,9-dihydro-9-oxo-3-phenyl-1,2-dimethyl seter (CA INDEX NAME) (Continued)

858249-88-4P, 5H-Furo[3,2-b]xanthene-6,7-dicarboxylic anhydride,
5a,6,7,8-tetrahydro-4-methoxy-5-oxo-8-phenyl- 858249-90-8P,
5H-Furo[3,2-b]xanthene-6,7-dicarboxylic anhydride,
5a,6,7,8-tetrahydro-4,11-dimethoxy-5-oxo-8-phenyl- 859780-23-7P,
1,2-Xanthenedicarboxylic acid, 1,2,3,9a-tetrahydro-9-oxo-3-phenyl-,
1,2,3,9a-tetrahydro-9-oxo-3-phenyl1,2,3,9a-tetrahydro-9-oxo-3-phenylRiperper (Preparation)
(preparation of)
858249-88-4 CAPLUS
3H-Difuro[3,4-a12',3'-i]xanthene-1,3,12-trione,
3a,4,12a,12b-tetrahydro-11-methoxy-4-phenyl- (CA INDEX NAME)

858249-90-8 CAPLUS
3H-Diffuro[3,4-a:2',3'-i]xanthene-1,3,12-trione,
3a,4,12a,12b-tetrahydro-7,11-dimethoxy-4-phenyl- (CA INDEX NAME)

859780-23-7 CAPLUS
1H-Xanthene-1,2-dicarboxylic acid, 2,3,9,9a-tetrahydro-9-oxo-3-phenyl-,
1,2-dimethyl ester (CA INDEX NAME)

L16 ANSWER 238 OF 238 CAPLUS COPYRIGHT 2009 ACS ON STN
ACCESSION NUMBER: 1933:618 CAPLUS
DOCUMENT NUMBER: 27:618
ORIGINAL REFERENCE NO. 27:89h-i,90a-f
TITLE: Thiophenols. Thiochromanone and thioxanthone
AUTHOR(S): Bellavita, V.
SOURCE: CAZZETTA Chimica Italiana (1932), 62, 655-63
CODEN: GCITAP; ISSN: 0016-5603
DOCUMENT TYPE: Journal
LANGUAGE: Unavailable
GI For diagram(s), see printed CA Issue.
AB Thiophenols were used as a starting point for the preparation of compds.
containing
3 sulfurated nuclei, and the present work describes the preparation of thiochromanone (I) and thioxanthone derivs. of trithiophloroglucinol (II)
(the only known trithiophenol). Only a very small yield of II is obtained obtained

thea by the method of Pollak and Carniol (C. A. 3, 2955), whereas the yield is 60-70% if C6H3(SO2Cl)3 (5 g.) in EtOH (20 cc.) and powdered Sn (15 g.)

are heated 0.5 hr. at 50°, concentrated HCl (50 cc.) is added (keeping the temperature at 30°), the solution is refluxed 0.5 hr., steam-distilled

and the amorphous residue is crystallized from EtOH. Attempts were made to obtain the

In the asym. trithiophenol from o-C6H4(SO3H)2 to 3,4-(HO3S)2C6H3NH2, thence by diazotization to 3,4-(HO3S)2C6H3SO2H, and thence (with KMnO4) to 1,3,4-C6H3(SO3H)3. The latter is probably formed in alkaline medium,

identify it, it has to be transformed into a sulfonyl chloride, amide or other derivative, and by the action of PC15 there is a simultaneous

of the sulfurated groups into sym. position, so that the only product is sym-C6H3(SO2C1)3. The aqueous Na salt of II and C1CH2CH2CO2H, heated (until a

ll a little acidified with HCl gives no mercaptan odor), filtered, acidified with H2SO4 or HCl, and the precipitate recrystd. from boiling water,

vield

trithiophloroglucinolpropionic acid, 1,3,5-C6H3(SCH2CH2CO2H)3 (IIII), m. 171-2°. III in concentrated H2SO4, heated 0.5 hr. at 50-60° (SO2 is evolved and the solution turns dark orange-red), poured into

the precipitate (a mixture of products) washed with water and dilute Na2CO3, and

03, and recrystd. from boiling AcOH or EtOH, yields I, orange-yellow, does not fuse up to 320°. The alkaline wash liquor from the preparation of I contains a mixture of 2 intermediate less dehydrated products, which can

be separated by acidifying with H2SO4, washing the precipitate with cold water, boiling the residue in water (1 product is much more soluble than the other), filtering hot, and cooling the filtrate, which yields, after further purification of the precipitate with boiling water, monothiochromanone-3,5-dithiopropionic acid (TV), light yellow, m. 216°. The residue insol. in boiling water, purified from EtOH, yields dithiochromanone-5-thiopropionic acid (V), golden yellow, m. 224-5°. II (5 g.) in aqueous NaOH (13 g.) and diazotized anthranilic acid (from 12 g.), heated until no more N is evolved (the solution turns orange-red), filtered, dilute H2SO4 added, the precipitate washed with hot AcOH,

L16 ANSWER 238 OF 238 CAPLUS COPYRIGHT 2009 ACS on STN (Continued) dissolved in aq. Na2CO3, repptd. with H2SO4 and recrystd. from EtOH or AcOH, yield benzene-1,2,3-rithio-o-benzoic acid, 1,3,5-C683(SC6H4CO2H-O)3 (VI), flesh-red, m. 300°. VI and concd. H2SO4, heated 6-7 hrs. on a water bath (the soln. turns an intense orange-red), poured into ice water, the ppt. washed with water, aq. Na2CO3, and water (its insoly, precludes its crystn. from any solvent), yield trithioxanthone (VII), amorphous, orange-red, remains unaltered up to 320°. The alk. wash liquor from the prepn. of VII acidified with H2SO4, the ppt. washed with water, and crystd. from AcOH by concn., yields monothioxanthone-3,5-dithio-o-benzoic acid (VIII), orange-yellow, turns slightly brown around 300°, but does not change further up to 320°.

IT 858846-56-3P, Thioxanthone, 1,3-bis(o-carboxyphenyl)RL: PREP (Preparation)
(preparation of)
RN 858848-56-3 CAPLUS
CN Benzoic acid, 2-[1-(2-carboxyphenyl)-9-oxo-9H-thioxanthen-3-yl]- (CA INDEX NAME)



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